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**Genomics of invasion: Diversity and selection in introduced populations
of monkeyflowers (*Mimulus guttatus*)**

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ABSTRACT

Global trade and travel is irreversibly changing the distribution of species around the world. Because introduced species experience drastic demographic events during colonisation, and often face novel environmental challenges from their native range, introduced populations may undergo rapid evolutionary change. Genomic studies provide the opportunity to investigate the extent to which demographic, historical, and selective processes shape the genomic structure of introduced populations by analysing the signature that these processes leave on genomic variation. Here we use next-generation sequencing to compare genome-wide relationships and patterns of diversity in native and introduced populations of the yellow monkeyflower (*Mimulus guttatus*). Genome resequencing data from ten introduced populations from the United Kingdom (UK) and 12 native *M. guttatus* populations in North America (NA), demonstrated reduced neutral genetic diversity in the introduced range, and showed that UK populations are derived from a geographic region around the North Pacific. A selective-sweep analysis revealed site frequency changes consistent with selection on 5 of 14 chromosomes, with genes in these regions showing reduced silent site diversity. While the target of selection is unknown, genes associated with flowering time and biotic and abiotic stresses were identified within the swept regions. The future identification of the specific source of origin of introduced UK populations will help determining if the observed selective sweeps can be traced to un-sampled native populations or occurred since dispersal across the Atlantic. Our study demonstrates the general potential of genome-wide analyses to uncover a range of evolutionary processes affecting invasive populations.

41 **INTRODUCTION**

42 The introduction of species beyond their native ranges can affect ecological and evolutionary
43 interactions in the new habitat (Cox 2004; Phillips & Shine 2006; Liu & Pemberton 2009;
44 Ricciardi *et al.* 2013), and can negatively impact levels of local biodiversity, and result in high
45 economic costs (Pimentel 2002; Williams *et al.* 2010; Vila *et al.* 2011). Introduced populations
46 are often used models to investigate rapid genetic changes and adaptation to novel
47 environments, thus providing valuable insights into basic biological processes including local
48 adaptation (Sax *et al.* 2007; Prentis *et al.* 2008). In particular, genetic analyses continue to play a
49 central role in studies of the origin and establishment of introduced populations, as well as of
50 the mechanisms that permit the colonisation and drive the spread of populations beyond their
51 native range (Baker & Stebbins 1965; Lee 2002).

52 The genomic structure of non-native populations is influenced by a variety of processes
53 including population bottlenecks, multiple introductions, population expansion, gene flow
54 between populations, and selection, among others (Lee 2002). For instance, in populations
55 established after limited long distance dispersal events, the level of genetic diversity can be
56 significantly lower than in the native range, reflecting population bottlenecks (Lachmuth *et al.*
57 2010; Ness *et al.* 2012). However, introduction of multiple individuals from the same
58 population, or multiple introductions from genetically diverse source populations, can
59 counteract the loss of diversity or even result in higher levels of genetic variation within
60 introduced populations compared to native ones (Dlugosch & Parker 2008). The level of
61 standing variation in introduced populations is relevant to the colonisation process, as severe

bottlenecks and reduced diversity could indicate potential limitations for the rapid evolution of adaptive traits in novel environments (Barrett & Schluter 2008; Lachmuth *et al.* 2010; Siol *et al.* 2010; Messer & Petrov 2013). Severe bottlenecks resulting in globally reduced diversity may indicate that natural selection is mutation limited. However, genetic variation resulting from introduction of multiple individuals can provide ample standing variation for natural selection.

Genome wide studies have been employed to investigate genetic patterns in natural populations, including the relationship between native and introduced populations as well as invasion pathways of exotic plants and animals (Jahodová *et al.* 2007; Dlugosch *et al.* 2013; Tarnowska *et al.* 2013). Genome scans allow detecting selection acting on specific locations in the genome (Nielsen *et al.* 2005), and by comparing the sites under selection in the genomes of different populations, it is possible to identify candidates for genetic regions associated with local adaptation (Savolainen *et al.* 2013). A prerequisite to any genome wide study is identifying a large numbers of genetic markers, such as restriction site polymorphisms (e.g., AFLPs, Vos *et al.* 1995), or single nucleotide polymorphisms (SNPs). The growing access to high-throughput sequencing technologies at low costs opens the opportunity to conduct genome wide studies at an unprecedented depth, even in non-model organisms (Prentis *et al.* 2010; Twyford & Ennos 2012; Ellegren 2014).

The generation of genome wide markers by high throughput sequencing can employ methods for genome complexity reduction such as transcriptome sequencing (Dlugosch *et al.* 2013) or RAD sequencing (Davey *et al.* 2011; Roda *et al.* 2013). However, for the increasing number of species in which a reference genome is available, whole genome re-sequencing

allows genotyping markers, such as SNPs, which may occur at high densities across the genome (Davey *et al.* 2011; Twyford & Ennos 2012; Savolainen *et al.* 2013). Importantly, whole genome resequencing removes many of the ascertainment biases associated with SNP chips or other genome reduction technologies. The dense marker saturation achieved through genome resequencing is particularly useful for detecting the footprint of selection acting on specific locations across the genome. For instance, selective sweeps, in which selection drives previously rare alleles to fixation, also reduces diversity at neighbouring regions around the selected site (Messer & Petrov 2013). The signal left behind by selective sweeps can be detected by comparing patterns of variation along the genome with the level expected under a null model. Hard selective sweeps, where a single variant is driven to fixation, leave a characteristic footprint in the genome, which can be identified using summary statistics such Tajima's *D* or the composite likelihood ratio (CLR) (Nielsen *et al.* 2005; Messer & Petrov 2013). These statistics may be particularly powerful to detect recent selective sweeps as linkage disequilibrium (LD) between the selected site and the surrounding variation is expected to be highest immediately following the fixation of the adaptive allele.

Genomic studies of native and introduced populations can uncover demographic, historical, and selective processes by analysing the signature that these processes leave on genomic variation. Here we use whole genome re-sequencing to assess the relationship between native and introduced populations, and to uncover selective episodes in specific regions of the genome of introduced populations. We study the yellow monkeyflower (*Mimulus guttatus*, Phrymaceae), a species that has long been used as a model for ecological and evolutionary

104 studies in its native range (Vickery 1959; Wu *et al.* 2008), and which has become naturalised in
105 Eastern North American, New Zealand, Iceland, the Faroe Islands, and Western Europe (van
106 Kleunen & Fischer 2008; Murren *et al.* 2009; Tokarska-Guzik & Dajdok 2010), becoming
107 particularly widespread in the United Kingdom (Vallejo-Marín & Lye 2013). *Mimulus guttatus* is
108 ideally suited for studying the ecological genomics of non-native populations due to its recent
109 introduction and spread (<200 years), abundant information on the ecology and evolution of
110 native populations, and the availability of a full genome sequence, which provides a backbone
111 for analysing and interpreting patterns of genetic variation in introduced populations. The
112 relatively small genome of *M. guttatus* (1N = 430 MB), makes this species a good candidate for
113 population genomic studies through re-sequencing, as multiple individuals can be analysed with
114 a relatively small budget. We analysed previously available and newly generated whole genome
115 sequence data for 12 native and 10 introduced British populations of *M. guttatus*, as well as five
116 additional related taxa ($n = 35$ *Mimulus* genomes in total). Our data set allowed us to address
117 three specific aims: (1) to determine the level of genome-wide diversity present in introduced
118 populations of *M. guttatus* in the United Kingdom; (2) to investigate the genetic relationships of
119 native and introduced populations; and (3) to search for evidence of hard selective sweeps in
120 introduced populations.

METHODS

Study system

The *Mimulus guttatus* species complex includes a set of phenotypically variable, interfertile taxa with a native range of distribution in Western North America from northern Mexico to Alaska (Grant 1924; Wu *et al.* 2007). Within this complex, populations of *M. guttatus* Fischer ex DC. (Grant 1924) show marked variation in characteristics including life history (annual/perennial (Hall & Willis 2006; Lowry & Willis 2010), mating system (Ritland 1990; Dole 1992), phenology (Hall & Willis 2006; Friedman & Willis 2013), floral morphology (Fishman *et al.* 2002), edaphic adaptations (e.g. tolerance to elevated concentrations of heavy metals or salt, Macnair & Watkins 1983; Lowry *et al.* 2008; Lowry *et al.* 2009), habitat preferences (Wu *et al.* 2008), chromosome number (most populations are diploid: $2n = 2x = 28$, but tetraploids also occur in the native range, Sweigart *et al.* 2008), and clonal growth (Dole 1992; van Kleunen 2007), among others. This incredible diversity has led some taxonomists to subdivide *M. guttatus* into numerous morphological species (e.g., Pennell 1951; Nesom 2012). Here we adhere to the broader circumscription of *M. guttatus* Fischer ex DC. (Grant 1924; Wu *et al.* 2008).

Mimulus guttatus was introduced into the British Isles in 1812, and the first naturalised populations were reported in England around 1830 (Roberts 1964; Parker 1975). In the UK, *M. guttatus* is currently widespread and occurs in wet habitats along the banks of rivers and streams, in ditches, marshy areas, and other wet places (Stace 2010; Vallejo-Marín & Lye 2013). It propagates via both seeds and clonally through lateral stems that root freely at the nodes.

The source of the first naturalised populations of *M. guttatus* in the UK is unknown, but one of the earliest specimens of this taxon to reach Europe was derived from material collected by Langsdorff between 1806 and 1810 in the Aleutian Islands in Alaska, and transmitted to the Botanic Gardens at Cambridge (Sims 1812; Pennell 1935, p. 116). The use of *Mimulus spp.* as a horticultural species in Victorian England, as reflected by being readily available in seed catalogues of the time (e.g., Gardeners' Chronicle 1852), raises the distinct possibility that *M. guttatus* was introduced into the UK on repeated occasions and from multiple sources.

Population sampling

Analysing genomes across a wide geographic scale represents a trade-off between the numbers of individuals vs. populations sampled. The goal of this study was to determine the introduction history of *Mimulus* into the UK, the effects of the introduction on nucleotide diversity, and to identify signals of selective sweeps that are common across the UK. To do this, we sought to obtain samples from geographic disparate regions from across the UK. Obtaining geographically distant samples increases the likelihood of identifying introductions from multiple different donor populations. This sampling strategy also facilitated our goal of identifying selective sweeps shared across the UK *M. guttatus* populations. Population specific selective sweeps caused by local adaptation to narrow geographic and ecological niches in the UK are not detected in our analyses and would require multiple individuals from the specific population of interest. Previous molecular analyses of *Mimulus guttatus* have demonstrated that a scattered sampling design, with 1 individual per population, is sufficient to capture regional differentiation, and can avoid clustering biases resulting from sampling multiple individuals from

fewer populations (Oneal *et al.* 2014). In total, we analysed genome data from 27 populations: 12 *M. guttatus* populations in the native range, 10 UK *M. guttatus*, and five outgroups. From one of the native populations (Iron Mountain, IM) we sampled an additional 8 individuals, which allowed us to explore the sensitivity of our findings to the particular individual sampled within a population.

Introduced populations

We sampled 10 populations of *M. guttatus* spanning the range of distribution of this species in the British Isles (Table 1; Fig. 1). The northernmost population came from the Shetland Islands (QUA, N 60.105° W 1.227°) and the southernmost from Cornwall, England (CRO, N 50.163°, W 5.293°). A population from Northern Ireland was also included (VIC, N 54.763°, W 7.454°). Non-native populations were collected from banks of canals, streams or rivers (HOU, CER, VIC, AYR, DBL, TOM, and PAC), on roadside ditches (QUA), on waterlogged ground in an abandoned field (CRO), or in a bog near a small stream (TRE). A single wild-collected individual per population was randomly selected from each population for sequencing.

Native range populations and outgroups

We obtained sequence data from the Sequence Read Archive (SRA) (<http://www.ncbi.nlm.nih.gov/sra>) from 12 native populations of *M. guttatus*, and five outgroups within section Simiolus: *M. nasutus* (SF), *M. cupriphillus* (MCN), *M. platycalyx* (CVP), *M. micranthus* (EBR), and *M. dentilobus* (DENT). The 12 native populations of *M. guttatus* covered a linear transect of approximately 2800km from Haida Gwaii (Queen Charlotte Islands), British Columbia (TSG, N 53.419°, W 131.916°) to Arizona (PED, N 32.711°, W 110.628°) (Table

1). Populations in the native range occurred in a diversity of wet habitats including river and stream banks, seeps, beach dunes, bogs, and springs. A single individual represented all but one of the native populations. In the case of the Iron Mountain population (IM), we were able to obtain data for nine separate individuals. Mean coverage per genotyped base per individual ranged between 4 – 29x with an average of 10x.

DNA isolation and sequencing

We collected leaf tissue of British *M. guttatus* individuals (one per population) in the field and preserved it in re-sealable plastic bags with self-indicating silica gel (Fisher Scientific, Loughborough, UK) at room temperature. This dry tissue was used for DNA extraction using the Leaf MasterPure total DNA extraction kit (Cambio Ltd, Cambridge, UK). DNA libraries were created and barcoded using the Nextera DNA sample preparation kit (Illumina, San Diego, California), which uses a transposon based method to randomly tag DNA for multiplexed sequencing. After library construction, an Agilent Bioanalyzer (Santa Clara, California) was used to measure length distribution of library, and a fluorometer (Qubit 2.0, Life Technologies, Paisley, UK) was used to measure concentration. Equimolar quantities of each library were pooled and sequenced in an Illumina HiSeq 2500 rapid-run producing 150 base-pair paired-end reads. Overall, we obtained raw coverage of 1.5 – 11x per individual with an average of 5.7x. Raw sequence data for UK *Mimulus* samples is deposited in the JGI SRA (SRA accession numbers will be available upon acceptance of the manuscript).

Sequence data analysis

Genome alignment and SNP genotyping

Raw reads were aligned to the *Mimulus guttatus* v2.0 genome available from Phytozome (<http://www.phytozome.net>) using *bowtie2* (Langmead & Salzberg 2014) using `–fast-local` allowing soft-clipping of poorly mapped read ends. After alignment, *Picard tools* (<http://picard.sourceforge.net>) was used to remove duplicates, add read groups, and verify that all mate information was accurate. After processing in *Picard tools*, the *Genome Analysis Toolkit* (GATK, DePristo *et al.* 2011) was used to call genotypes using the 'Unified Genotyper'. Minimum alignment quality was 25 and base quality was 25. Called genotypes were filtered to include genotypes with a call quality threshold of Q30 or greater. Insertions, deletions, and heterozygous sites were not included in subsequent analyses. Detailed command-line methods can be found in the Supplementary Information. After all filtering, mean coverage per genotyped base per individual for the 10 UK samples ranged from 1.7 – 5.8x with an average of 3.5x . Of the 293 Mb located on the main 14 genomic scaffolds (representing 14 linkage groups), after all filtering, 71 Mb were genotyped in at least one of the UK individuals. A total of 18.3, 18.5, and 8.9 Mb were genotyped in 8, 9, or 10 of the UK samples, respectively (Fig. S7).

Measures of genetic diversity

Nucleotide diversity at silent and non-silent sites was calculated using software described in Zhang *et al.* (2006). Briefly, genomes for all sequenced lines were recalled using the genotype data. Missing data was not imputed. Measurements of pairwise synonymous (π_{syn}) and non-

synonymous nucleotide diversity ($\pi_{\text{non-syn}}$) were calculated through pairwise comparison of coding sequences. Coding sequences were extracted from the recalled genomes using the gff3 gene annotation file available on phytozome.net. A Fisher p-value associated with each diversity value and indicates the confidence of that particular value. We only considered π_{syn} and $\pi_{\text{non-syn}}$ values for genes with a Fisher p-value ≤ 0.001 and alignment length greater than 200 bases. In addition to calculating nucleotide diversity at synonymous and non-synonymous sites, whole genome alignments were used to calculate genome-wide nucleotide diversity (π) in sliding windows using *VariScan* (Hutter *et al.* 2006). Windows of 50,000 genotyped bases and overlapping steps of 1000 bases were used.

Genetic relationships between introduced and native *M. guttatus*

In order to determine the genetic relationships between introduced and native populations we conducted an analysis of genetic similarity using a random subset of 1,400,000 SNPs. To create this data set, we randomly selected 100,000 SNPs for each of the 14 major linkage groups (chromosomes) that were genotyped in at least 30 individuals (out of 35). Our SNP data set is therefore not subject to ascertainment bias arising from selecting, for example, only coding or non-coding SNPs (Garvin *et al.* 2010). Instead the SNP dataset analysed here should represent a snapshot of the total genetic diversity of each sample and be shaped by both neutral and non-neutral processes (Helyar *et al.* 2011). Within each linkage group, neighbouring SNPs were separated by 209 bp on average (209 ± 3.34 ; mean \pm SE). Each SNP was coded as “0” if it matched the reference allele, and “1” for the alternative allele. In this analysis we included all native and introduced individuals, and the five outgroups ($n = 35$ individuals). Multiple

individuals from IM were included as a reference of the variation seen within a single population.

We constructed a genetic distance matrix using p -distance (the proportion of nucleotide sites that differ between a pair of sequences) from the binary SNP data using the package *ape* (Paradis *et al.* 2004) in *R* ver. 3.0.3 (R Development Core Team 2014). The combined distance matrix was then used to estimate the relationships between all samples using a neighbour joining (NJ) analysis in *ape*. Support for nodes in the NJ tree was calculated using 100 bootstrap replicates. Trees were drawn using *FigTree* v. 1.4.0 (Rambaut 2014). The NJ distance-based approach used here is appropriate for genome-wide analyses (e.g. Brandvain *et al.* 2014), as maximum likelihood and Bayesian phylogenetic methods depend on specifying a mutational model, which is not practical for genome-wide data. We also conducted a Principal Component Analysis (PCA) using the function *glPca* in *adeigenet* (Jombart & Ahmed 2011). This analysis provides an independent estimate of the relationships between native and introduced populations, and can be used to compare with the results of the NJ analysis. For the PCA, we selected only one individual for each of the 12 native and 10 introduced populations of *M. guttatus*. The identity of the particular individual chosen from the IM population had no qualitative effect on the relationships inferred from the PCA (Fig S8), and similarly, randomly choosing one IM individual instead of nine for the NJ analysis did not change the tree topology (data not shown).

Selective sweep analysis

Regions in the genome showing the signature of selective sweeps were detected using the parametric approach described in Nielsen *et al.* (2005), and implemented in the program *SweepFinder*. This method compares the observed site frequency spectrum within local regions in the genome (windows) against the background site frequency spectrum seen across the entire genome (or linkage group), and calculates statistical departures from this background expectation using a composite likelihood ratio (CLR). Importantly, the null hypothesis employed by this method is derived from the background data itself and does not depend on specific population genetic models or assumptions about demographic equilibrium (Nielsen *et al.* 2005), which are unlikely to hold in recently introduced populations. *SweepFinder* is robust to models that include population growth with recombination (Nielsen *et al.* 2005).

One potential issue with *SweepFinder* is that it is sensitive to SNP density (Nielsen *et al.* 2005). To account for both shared ancestral sweeps and artefacts due to genotype density, we independently analysed the North American (NA) and UK data using the exact same criteria. Ten samples from the North American populations (AHQ, BOG, DUN, IM, LMC, MAR, PED, SWB, TSG, YJS) were chosen based on the results of the genome-wide relationship analysis. Next, independently, for both the NA and UK datasets, we determined whether a given site was polymorphic and asked how many individuals were genotyped at that particular site. Using this information, we choose sites that were genotyped in at least 8 individuals (of the 10 total) in both the UK and NA samples and were polymorphic in at least one of these populations. Thus, we ended up with a dataset that included the exact same number of sites from the exact same

genomic locations. Next, we calculated genome-wide SFS for each dataset (UK and NA). Each chromosome was divided into 5000 bins and the SFS within each bin was compared to the genome-wide SFS to look for signals of a selective sweep using the parametric approach described (Nielsen *et al.* 2005) and implemented in *SweepFinder*.

After running *SweepFinder*, we independently plotted the genome-wide CLR distribution for UK and NA samples. Within the NA samples, all genomic locations with CLR scores above the median value were marked for masking. Liberal masking based on the NA analyses removes hard sweeps that occurred in the last common ancestor, and removes artefacts due to variable genotype density. The NA *SweepFinder* results were used to mask the UK genome *SweepFinder* results. Only genomic positions that survived masking were considered in further analyses. Within the UK, the top 1% CLR outliers were identified as regions that have possibly experienced a hard sweep and subjected to further investigation. Gene coordinates are available in the gff3 gene annotation on Phytozome and genes with positions at least partially overlapping the swept regions were extracted for further analyses.

RESULTS

Nucleotide diversity in *M. guttatus*

Overall genome-wide nucleotide diversity in the UK was $\pi = 0.015$ (Fig. S1). Figures S2 and S3 show patterns of nucleotide diversity across the genome for both native and introduced populations. For genes, within UK samples, diversity at silent sites was $\pi_{\text{syn}} = 0.0325$ while non-synonymous diversity was $\pi_{\text{non-syn}} = 0.0035$ (Fig. S4). Within the native North American

populations, nucleotide diversity was calculated through comparisons of 10 individuals (same 10 individuals used as the NA samples in sweep analyses). Genome-wide diversity within the NA samples is $\pi = 0.031$ (Fig. S1). Synonymous diversity within NA is $\pi_{\text{syn}} = 0.0610$ while non-synonymous diversity is $\pi_{\text{non-syn}} = 0.0075$ (Fig. S4). Comparing NA and UK nucleotide diversity indicates an overall reduction in the introduced populations of approximately 50%.

Genetic relationships of native and introduced populations

The relationships between 22 native and introduced populations of *Mimulus guttatus*, and five outgroups based on the genetic distance of 1,400,000 SNPs distributed across the genome is shown in Fig. 2. All the population level nodes in this NJ tree had a bootstrap support of 100%. The NJ tree shows that all 10 UK populations form a single well-supported clade (Fig. 2). The UK clade is most closely related to the native TSG population, a coastal perennial *M. guttatus* from Graham Island (Queen Charlotte Islands) in British Columbia (Lowry & Willis 2010). The UK and TSG samples form part of a clade of populations located north of the N 40° parallel, and which includes other inland perennial (BOG and YJS), and annual populations (AHQ, IM, and MAR) (Figs. 1 and 2; Table 1). The NJ tree shows a second clade composed mainly of more southern populations, and which includes inland plants (LMC, REM), coastal perennials (SWB, DUN), as well as two annual outgroups (CVP, *M. platycalyx*; EBR, *M. micranthus*). The DUN population is the only one in this group located north of the N 40° parallel (Fig. 1). Finally, the NJ tree shows a third clade including three of outgroups (MCN, *M. cupriphilus*; SF, *M. nasutus*; DENT, *M. dentilobus*), as well as the two most southern populations of *M. guttatus*, MED, an annual inland population, and PED, an inland perennial from Arizona. Our results indicate that native

M. guttatus is separated into two main clades corresponding mostly to geographic location (North and South groups), and not to different life histories (annual/perennial) or habitat types (coastal/inland), as has been recently described by Brandvain *et al.* (2014). Finally, our NJ analysis also indicates that *M. cupriphilus* and *M. platycalyx* are nested within broadly circumscribed *M. guttatus* (Fig. 2).

The results of the PCA show clear support for a close relationship between all UK samples, and also indicate that the most genetically similar native population sampled here is the coastal perennial TSG (Fig. 3). The first principal component separates the North and South groups of native *M. guttatus*, with DUN partly overlapping with the North group. The second principal component separates the UK samples from most of the other northern accessions (Fig. 3). Together, the NJ and PCA show a common ancestry of British *Mimulus guttatus*, and its association with northern, native populations.

Evidence of selection in introduced populations

Our analysis of selective sweeps in native and introduced populations identified several genomic regions displaying changes in the site frequency spectrum (measured using the CLR statistic), consistent with the signatures of positive selection acting in these regions (Fig. 4; results for all linkage groups are shown in Fig. S5). The comparison of high CLR regions in the separate analyses conducted in NA and UK samples, allowed us to detect selective sweeps shared by both native and introduced populations. By masking these high CLR sites in NA, we located genomic regions that are candidates for selective sweeps occurring after the separation of the clade leading to the UK populations, including potentially unsampled North American donor

populations. Moreover, the masking also accounted for possible regions with high CLR scores that were simply due to low genotyped SNP density. Our analysis revealed selective sweeps on five of the 14 linkage groups in the UK, which are not shared with NA samples (Fig. 4, Table 1). Regions with high CLR scores showed significantly reduced overall diversity: Diversity within sweeps was $\pi = 0.0076$, half that of nucleotide diversity outside sweeps $\pi = 0.0152$. ($p \leq 0.0001$; Figs. S6).

We identified a total of 299 genes located under the candidate region for selective sweeps within the UK clade (Table S1). Synonymous diversity for genes within sweeps was significantly lower than silent site diversity outside sweeps of ($\pi_{\text{syn}} = 0.0147$ vs. $\pi_{\text{syn}} = 0.0323$; $p \leq 0.0001$) (Fig. S6). Within sweeps, 28 genes had $\pi_{\text{syn}} < 0.01$ (Table S2) and only two genes under sweeps had silent site diversity (π_{syn}) above the genome-wide mean. Taking advantage of the annotated genome of *M. guttatus*, we recorded genes located within the swept regions, which included genes involved in flowering time, abiotic stress response including nutrient transport and freezer tolerance, and biotic stress responses (Table S1).

DISCUSSION

Our study represents the first genome resequencing study of native and introduced populations of *Mimulus guttatus*. By analysing whole genome sequences of 35 individuals from 22 populations we demonstrated that introduced plants in the UK are characterized by a ~50% reduction in synonymous (π_{syn}) genetic diversity, and that UK populations form a single clade, relative to the North American samples included here, suggesting a common origin for non-

native populations. Our analysis revealed changes in the site frequency spectrum at multiple locations across the genome, consistent with selective sweeps, some of which were restricted to UK samples. The reduction of both synonymous and non-synonymous nucleotide diversity in genes located within sweep regions, compared to genes outside of these sweeps, was consistent with the expected loss of genetic diversity in regions linked to selected loci. Future studies are required to assess whether selection is indeed responsible for the observed patterns of variation in the sweep candidate regions, as well as to determine if such selection acted before or after the introduction of *M. guttatus* into the UK. This study illustrates the potential of whole-genome sequencing studies to provide the initial steps for understanding the genomic consequences of invasion and rapid adaptation to new environments.

Origin and diversity of *Mimulus guttatus* populations in the UK

Determining the geographic origin of introduced populations provides a reference point for studies of the potential ecological and evolutionary changes occurring during the colonization and establishment phases of biological invasions (Milne & Abbott 2000). Our sample of North American populations includes a large part of the native range of *M. guttatus* (Fig. 1), but admittedly still represents a small fraction of populations from this widely distributed taxon (Grant 1924). However, we were able to include populations with different life histories, morphologies, and habitat preferences (Table 1), including different ecological and morphological groups (Lowry *et al.* 2008; Lowry & Willis 2010).

Our analysis of genome-wide polymorphism revealed two main clades of *M. guttatus* in the native range, which broadly correspond to their geographic origin (North and South groups;

Brandvain *et al.* 2014). An exception to the geographic arrangement of these two groups of *M. guttatus* is the DUN population. Although geographically located in the northern range (Fig. 1), DUN is nested within the southern group (Fig. 2), and may represent a secondary dispersal or reflect an introgression event between these two groups (Brandvain *et al.* 2014). Interestingly, the North and South clades of *M. guttatus* include populations with contrasting habitats, morphologies, and life histories (Table 1), suggesting that taxonomic groupings based on general morphological and ecological attributes are unlikely to correspond to monophyletic clades (Nesom 2012). At the same time, the polyphyletic nature of the *M. guttatus* species complex is reflected in our results by the fact that a *M. guttatus* population (MED) is nested within outgroup taxa (SF: *M. nasutus*, and MCN: *M. cupriphilus*; Fig. 2), while *M. platycalyx* (CVP) and *M. micranthus* (EBR) fall within *M. guttatus* populations. Elucidating the phylogenetic relationships within the *M. guttatus* species complex has proven challenging (Beardsley *et al.* 2003; Beardsley *et al.* 2004), but the use of genome-wide sequences in a phylogenetic context (Wagner *et al.* 2013) could provide a tool to establish the genetic relationships among populations of this interfertile group.

The genetic structure observed in indigenous populations, allowed us to determine with confidence that introduced populations in the UK are most genetically similar to populations from the northern end of the native distribution. In particular, our results indicate that UK samples are most genetically similar to the coastal perennial population of TSG (Figs. 2 and 3). However, without additional sampling, the exact source for introduced populations in the UK remains unknown. The genetic similarity within UK populations, which fall within a single clade

(Fig. 2), suggests that this part of the introduced range has been established either via a single introduction event or, perhaps more likely, via multiple introductions from closely related native populations. In order to determine the origin and number of introductions of UK *M. guttatus* with more accuracy, it will be necessary to conduct further sampling in the Alaskan end of the distribution, since historical records point to this region as a potential source for the first specimens of this taxon in Europe (Sims 1812; Pennell 1935, p. 116).

Despite a 50% reduction in nucleotide diversity (π) in non-native *M. guttatus*, UK populations still harbour a significant amount of diversity among populations. In particular, the estimate of nucleotide diversity at synonymous sites in UK populations ($\pi_{\text{syn}} = 0.0325$) is above the median estimate of neutral nucleotide diversity for outcrossing flowering plants ($\pi = 0.0148$), and much higher than the estimate for selfing taxa ($\pi = 0.0035$) (Leffler *et al.* 2012). The diversity observed in UK populations suggests that the bottleneck experienced by *M. guttatus* during invasion was weak enough to permit the maintenance of a significant amount of nucleotide diversity. In general, introduced populations tend to show reduced genetic diversity compared to native ones, but phenomena including large initial propagule numbers, multiple introductions, and admixture can result in equal or higher levels of genetic variation (Dlugosch & Parker 2008). The existence of non-synonymous variation in UK *M. guttatus* ($\pi_{\text{non-syn}} = 0.0035$) is potentially important for naturalisation and spread in the new range, as introduced populations may be able to respond to new selective pressures from standing genetic variation, and are not necessarily limited by mutation rate (Prentis *et al.* 2008). Adaptation from standing

genetic variation is particularly important for invasive species as it may allow responding more rapidly to novel selective pressures (Barrett & Schluter 2008).

Detecting selection in non-equilibrium populations

Our study uncovered several areas of the genome that bear the signature of positive selection.

The signal left by selective sweeps include increased frequency of previously rare derived alleles, reduction of variation in linked sites, and increase in local linkage disequilibrium (Smith & Haigh 1974; Barton 1998). However, past demographic changes can confound the pattern of a selective sweep (Barton 1998; Barton & Etheridge 2004). This is particularly the case in populations that have experienced a bottleneck and subsequent population growth with recombination (Barton & Etheridge 2004; Nielsen *et al.* 2005). Furthermore, recombination between lineages that either escape or not the population bottleneck will result in a combination of short and long coalescent branches creating a sweep like pattern in the SFS (Pavlidis *et al.* 2010). Thus, inferences of selection based on genome scans in non-equilibrium populations must be done with caution.

The implementation that we used to detect the signature of positive selection (*SweepFinder*), is relatively robust to the effects of bottlenecks and population expansion, and is particularly powerful to detect recent selective sweeps (Nielsen *et al.* 2005; Pavlidis *et al.* 2010). Recent studies have shown that *SweepFinder* performs better than other tests, such as the ω -statistic, in non-equilibrium populations (Pavlidis *et al.* 2010). We consider that the very recent introduction of *M. guttatus* into the UK, relatively weak population bottleneck, and lack of within-UK population structure, make *SweepFinder* an appropriate method for detecting recent

and strong sweeps in the introduced range. Moreover, the independently analysed NA data using the exact same SNP sampling strategy allowed us to mask shared sweeps and eliminate false positives due solely to SNP sampling and variable site density. Therefore, we consider that the genomic regions identified here are strong candidates for selective sweeps in the lineage leading to UK populations. Nevertheless, it is important to recognise that we cannot currently determine whether the selection events happened in an ancestral (unsampled) native population or after the dispersal of *M. guttatus* to the UK. Establishing the timing of the potential selective event is essential to determine if introduced populations can exploit novel environments through (pre-)adaptations brought in from the native range, or whether adaptive evolution occurs subsequent to dispersal during the establishment and spread phases of a biological invasion (Maron *et al.* 2004; Colautti *et al.* 2009).

Genes within swept regions

The potential selective sweeps we detected in five *M. guttatus* chromosomes include approximately 300 genes. As predicted for selective sweeps, we found reduced diversity in both coding and non-coding genic regions under these sweeps (Fig S6). While it is possible to identify candidate regions for selective sweeps and to determine the genes located under these sweeps, it is not possible to know without direct testing which genes were the actual targets of selection. However, the selective sweeps identified here contain genes involved in flowering time, nutrient stress, and biotic stress (Table S2), and could be involved in adaptation to different day lengths, soil types, novel pathogens, and general responses to stress (e.g., Hodgins *et al.* 2012).

Of particular interest is the identification of selective sweeps in linkage group 8 (LG8). The selective sweeps at positions 2 Mbp (two million base pairs) and 5 Mbp of LG8 are exceptionally wide, consistent with very strong selection pulling a large haplotype block to high frequency very quickly giving very little time for recombination to break up linked sites. These regions of LG8 are gene rich suggesting that the width of the peak is not an artefact of reduced recombination associated with repetitive regions (Hellsten *et al.* 2013). Instead, these selective sweeps are located near or at a known inversion region of approximately 6Mb in length (Oneal *et al.* 2014). This inversion (*DIV1*), is polymorphic in the native range, and is associated with a number of morphological and life-history differences between annual and perennial ecotypes (Lowry and Willis 2010). Therefore this region is a good candidate for bearing adaptive variation that could be selected for in the lineage leading to the introduced populations. Mapping experiments in *M. guttatus* have demonstrated that a large region on LG8 is involved in critical photoperiod to flower (Friedman & Willis 2013). The selective sweep we identified on LG8 is located near (within 200,000 base pairs) one of the major QTLs for critical photoperiod identified in Friedman and Willis' study. Flowering time is a crucial component of fitness in seasonal environments, and therefore it is expected to be under selection in the introduced range. Studies of invasive species have often demonstrated the potential for rapid evolution of phenology in the introduced range (Maron *et al.* 2004; Colautti *et al.* 2009). The geographic distribution of introduced populations at high latitudes (approximately between 50° N and 60° N in the UK), suggests that these populations experience day length conditions similar to the northern end of the distribution of *M. guttatus* in North America. It would be of interest to

determine if genes involved in the control of flowering under long days are under selection in northern indigenous or in introduced populations. One such candidate is *Migut.D02071*, a phytochrome-associated protein phosphatase gene located in one of the selective sweep regions in LG4 (Table S2). In *Arabidopsis thaliana*, a similar gene (*ATFYPP3*) participates in the regulation of flowering time in long days (www.string-db.org). Preliminary data suggests that UK populations require long days (16hr) to flower (Vallejo-Marín, unpublished). Future studies could address whether flowering time has indeed played a role in the establishment of introduced populations in high latitudes.

To conclude, our results demonstrate the enormous potential for whole-genome sequencing studies to contribute to the study of non-native populations. With a modest sequencing effort we were able to quickly obtain previously unavailable information on the origin and diversity of introduced populations of *M. guttatus*, as well as on the genomic consequences of biological introductions, including identifying potential regions under selection. As whole genome reference data becomes available for other non-model organisms, resequencing studies are likely to be increasingly used to study the history and consequences of biological invasions, and to establish the contribution of adaptive processes to shaping the genomes of rapidly evolving populations.

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703 **Data Accessibility**

704 **DNA sequences:** NCBI SRA (individual population codes in parentheses): SRR1462346 (10-AYR-10), SRR1475232 (10-CER-10),
705 SRR1475385 (10-DBL-20), SRR1481643 (10-HOU-17), SRR1481644 (10-QUA-47), SRR1482404 (10-TOM-23), SRR1482405 (12-
706 CRO-5), SRR1482406 (12-PAC-39), SRR1482407 (12-VIC-18), SRR1482409 (12-TRE-17).
707 **SNP Genotype data:** Dryad doi:10.5061/dryad.3gp32

708 **Author Contributions**

709 MVM and JP designed the research. MVM made the field surveys and collected the samples. JP and MVM organized laboratory
710 work. JP generated the genomic data and conducted the bioinformatic analyses. JP and MVM analysed the genomic data. MVM
711 and JP wrote the manuscript.

TABLES

Table 1. Populations sampled for genome resequencing of *Mimulus guttatus* and related taxa. A single individual was sequenced per population, except for IM, where sequence data was available for nine individuals. Life history and, for native populations, their classification as coastal or inland, is provided for each population when available (Lowry *et al.* 2008; Lowry & Willis 2010).

Population	Taxon	Latitude	Longitude	Life history	Coastal/Inland	Location
<i>Mimulus guttatus</i> (Native)						
PED		32.711	-110.628	Perennial	Inland	San Pedro River, Pinal Co., AZ
MED		37.829	-120.345	Annual	Inland	Moccasin, Tuolumne Co., CA
REM		38.859	-122.410	--	Inland	Rumsey, Yolo Co., CA
LMC		38.864	-123.084	Annual	Inland	Yorkville, Mendocino Co., CA
SWB		39.036	-123.691	Perennial	Coastal	Sperm Whale Beach, Mendocino Co., CA
BOG		41.924	-118.804	Perennial	Inland	Bog Hot Springs, Humboldt, Co., NV
MAR		43.479	-123.294	Annual	Inland	Marshanne Landing , Douglas Co., OR
DUN		43.893	-124.130	Perennial	Coastal	Dunes, Lane Co., OR
IM		44.401	-122.151	Annual	Inland	Iron Mountain, Linn Co., OR
AHQ		44.431	-110.813	Perennial	Inland	Lonestar Basin Thermal Spring, Teton Co., WY
YJS		44.951	-114.585	Perennial	Inland	Yellowjacket creek, Lemhi Co., ID
TSG		53.419	-131.916	Perennial	Coastal	Graham Island, Haida Gwaii (Queen Charlotte Islands), British Columbia, Canada
<i>Mimulus guttatus</i> (Introduced)						
CRO		50.163	-5.293	Perennial	--	Crowan, Cornwall
TRE		50.498	-4.465	Perennial	--	Tremar Coombe, Cornwall, England
HOU		51.097	-1.508	Perennial	--	Houghton Lodge, Hampshire, England

CER		53.006	-3.549	Perennial	--	Cerrigydrudion, Denbigshire, Wales
VIC		54.763	-7.454	Perennial	--	Victoria Bridge, Northern Ireland
AYR		55.461	-4.625	Perennial	--	Ayr, Ayrshire, Scotland
DBL		56.197	-3.965	Perennial	--	Dunblane, Perthshire, Scotland
TOM		57.255	-3.368	Perennial	--	Tomintoul, Moray, Scotland
PAC		57.355	-3.336	Perennial	--	Packhorse Bridge, Speyside, Scotland
QUA		60.105	-1.227	Perennial	--	Quarff, Shetland Islands
Outgroups						
SF	<i>M. nasutus</i>	45.635	-120.914	Annual	Inland	Sherars Falls, Wasco Co., OR
MCN	<i>M. cupriphilus</i>	37.912	-120.724	Annual	Inland	McNulty Mine, Calaveras, Co., CA
CVP	<i>M. platycalyx</i>	38.372	-123.055	Annual	Inland/Coastal	Coleman Valley Road, Sonoma Co., CA
EBR	<i>M. micranthus</i>	39.631	-123.532	Annual	Inland	Branscomb, Mendocino Co., CA
DENT	<i>M. dentilobus</i>	NA	NA	--	--	NA

717

FIGURES

Figure 1. Location of the 22 *Mimulus guttatus* populations sampled in the native range in North America (left-hand side panel), and in the introduced range the United Kingdom (right-hand side panel). Notice the different scales in the two maps. The colour of the symbols corresponds to the clades shown in the neighbour joining tree in Fig. 2.

Figure 2. Relationships between native (North America) and introduced (United Kingdom) populations of *Mimulus guttatus* inferred from 1,400,000 SNPs, sampled at a density of 100,000 SNPs per chromosome. The neighbour joining tree was built from a matrix of pairwise genetic distance (p -distance) of 30 individuals of *M. guttatus*, four related taxa, and rooted with *M. dentilobus*. Branches leading to introduced populations are shown in purple. The two clades containing native *M. guttatus* in the North and South groups (see Results) are shown in blue and red, respectively. All nodes at the population level have a 100/100 bootstrap support. Population names as in Table 1.

Figure 3. Principal component analysis of twelve native (North America) and ten introduced (United Kingdom) populations of *Mimulus guttatus*. Introduced populations cluster in the lower left quadrat. Only one individual of the Iron Mountain (IM) population was included in this analysis. Colours denote values at first two principal component (PC) axes. Population names as in Table 1.

Figure 4. Evidence of hard selective sweeps within UK *M. guttatus* was found on 5 of the 14 linkage groups using *SweepFinder* (Nielsen *et al.* 2005). The figure shows selective sweeps after

738 masking for shared sweeps between native and introduced populations (see Methods). Dashed
739 line indicates genome-wide 1% outlier cutoff based on the composite likelihood ratio (CLR)
740 statistic. Only linkage groups with sweeps exceeding 1% outlier cutoff are shown here; CLR
741 profiles for all linkage groups are shown for both native and introduced populations in the
742 supplementary materials.

SUPPLEMENTARY MATERIAL

Figure S1. Distribution of genome-wide pairwise nucleotide diversity (i.e., the average number of nucleotide differences per site between two sequences, π) in native and introduced populations of the yellow monkeyflower, *Mimulus guttatus*. The left-hand side panel shows nucleotide diversity of 10 introduced populations in the United Kingdom, and the right-hand side panel shows nucleotide diversity for 10 native populations in North America (same 10 used in sweep analysis). Nucleotide diversity was calculated using windows of 50,000 genotyped base pairs (bp) and 1,000 bp overlapping steps.

Figure S2. Genome-wide pattern of pairwise nucleotide diversity (π) in 10 introduced populations of *Mimulus guttatus* in the United Kingdom. Each row represents one of the 14 major linkage groups (scaffolds). Nucleotide diversity was calculated using windows of 50,000 genotyped base pairs (bp) and 1,000 bp overlapping steps. The x-axis indicates the midpoint position of each window with respect to the *M. guttatus* v 2.0 reference genome (www.phytozome.org).

Figure S3. Genome-wide of pairwise nucleotide diversity (π) in 10 native populations of *Mimulus guttatus* in North America (AHQ, BOG, DUN, IM, LMC, MAR, PED, SWB, TSG, YJS). Each row represents one of the 14 major linkage groups (scaffolds). Nucleotide diversity was calculated using windows of 50,000 genotyped base pairs (bp) and 1,000 bp overlapping steps. The x-axis indicates the midpoint position of each window with respect to the *M. guttatus* v 2.0 reference

genome (www.phytozome.org).

Figure S4. Distribution of non-synonymous ($\pi_{\text{non-syn}}$) and synonymous (π_{syn}) pairwise nucleotide diversity in 10 introduced (left-hand side panels; United Kingdom), and in 10 native populations (right-hand side panels; North America) of *Mimulus guttatus*.

Figure S5. Selective sweeps identified in introduced populations (United Kingdom) of the yellow monkeyflower, *Mimulus guttatus*, across the 14 major linkage groups (scaffolds) in this species. The composite likelihood ratio (CLR) measures departures in the site frequency spectrum from background levels observed within linkage groups, and provide evidence of selective sweeps (Nielsen *et al.* 2005). Black dots indicate CLR scores above the 1% outlier cutoff, after masking (omitting) regions with significant CLR scores in North American samples. The x-axis indicates the position in base pairs with respect to the *M. guttatus* v 2.0 reference genome (www.phytozome.org). For full details of the analysis see Methods section.

Figure S6. Evidence for reduced pairwise nucleotide diversity (π and π_{syn}) in genes located within (sweep) vs. outside selective sweeps (no sweep) of introduced populations of *Mimulus guttatus* in the United Kingdom. The left-hand side panel shows pairwise nucleotide diversity across all sites within a gene (π), and the left-hand side panel shows pairwise nucleotide diversity in synonymous sites only (π_{syn}). For the left panel, nucleotide diversity was calculated using windows of 50,000 genotyped base pairs (bp) and 1,000 bp overlapping steps. Synonymous site nucleotide diversity was calculated for all genes within swept region. Boxplots show the median (horizontal line), 1st and 3rd quartiles (top and bottom of the box), and the 1.5

interquartile range (whiskers).

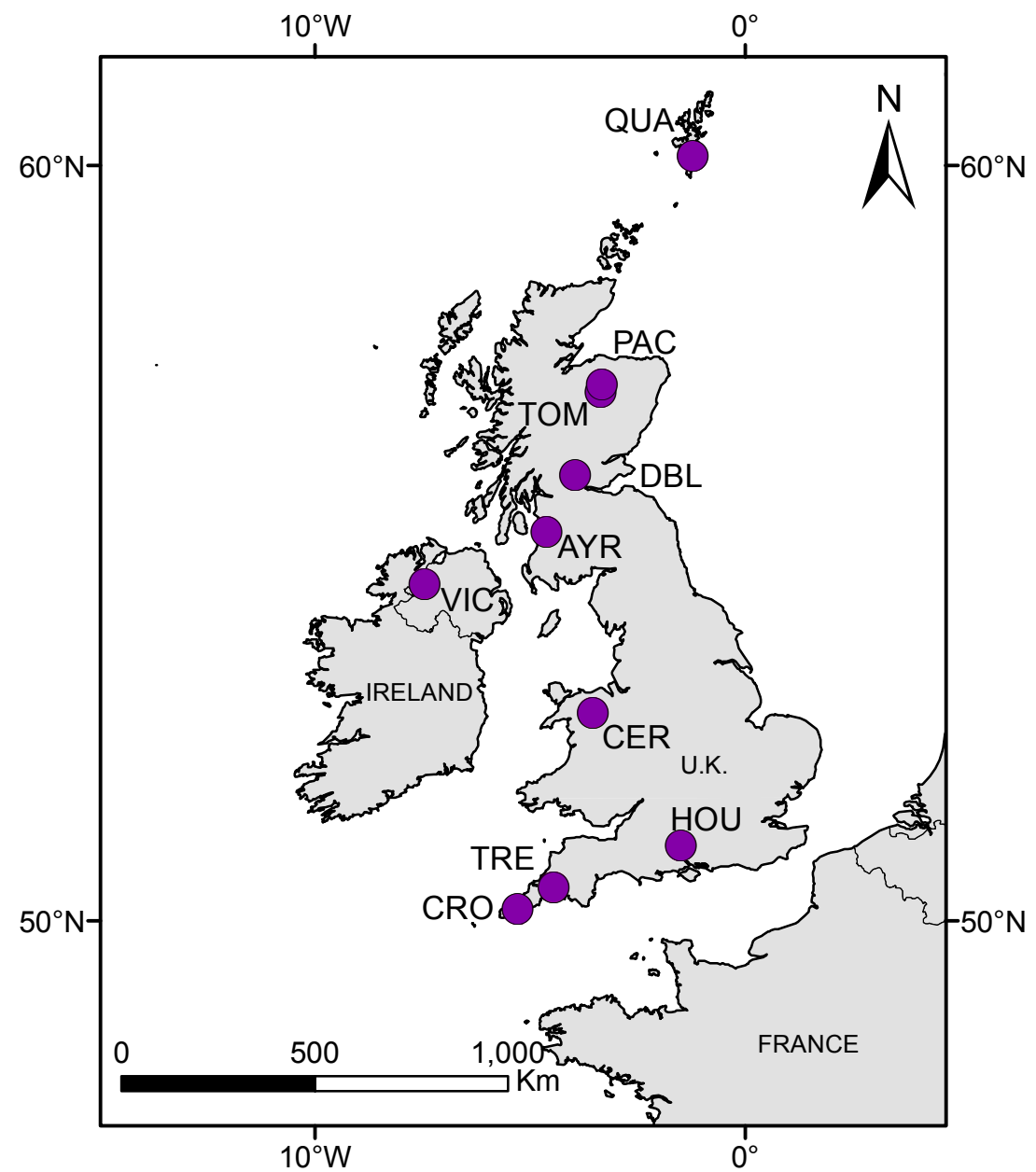
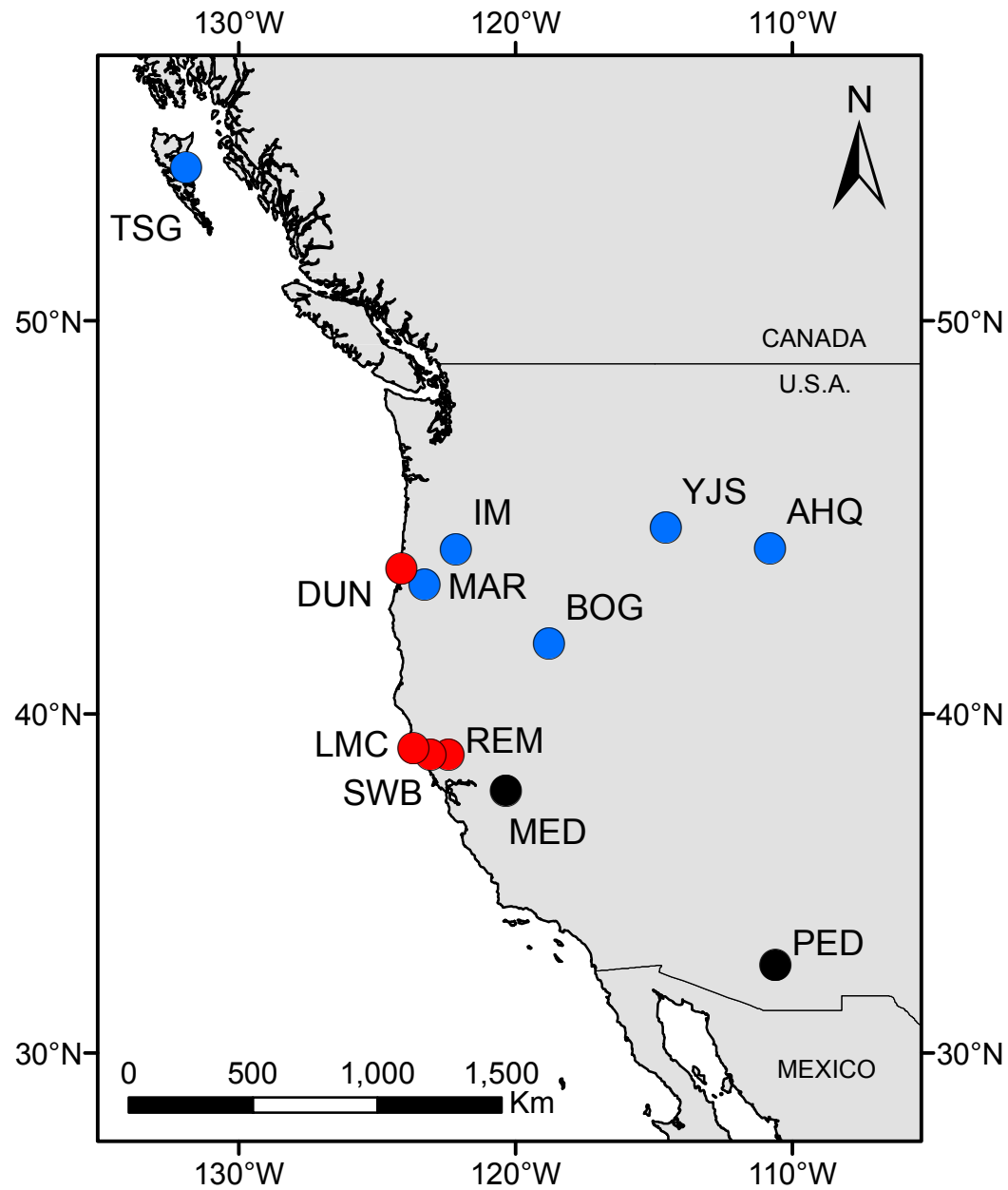
Figure S7. Genome-wide distribution of genotyped base pairs per individual across ten individuals from ten introduced populations of *Mimulus guttatus* in the United Kingdom (UK). The chart shows, for instance, that nine out of ten individuals were successfully genotyped at more than 18 million base pairs.

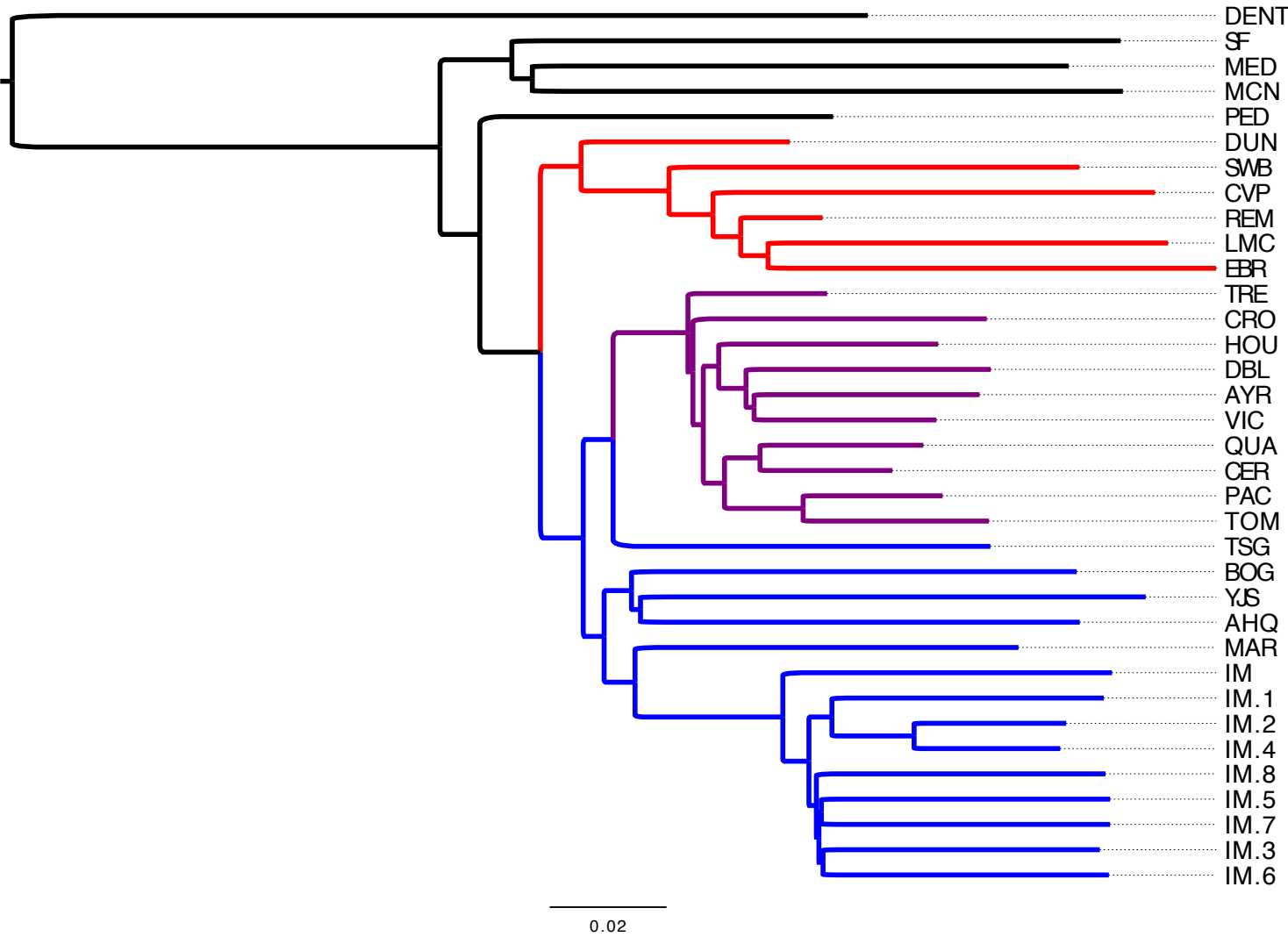
Figure S8. Principal component analysis (PCA) of 30 individuals of *Mimulus guttatus* from both native and introduced ranges, including multiple individuals of the Iron Mountain (IM) population. This PCA was based on 10,000 SNPs per linkage group for a total of 140,000 SNPs across the genome. Introduced populations fall in the bottom left quadrat. Population codes are defined in Table 1.

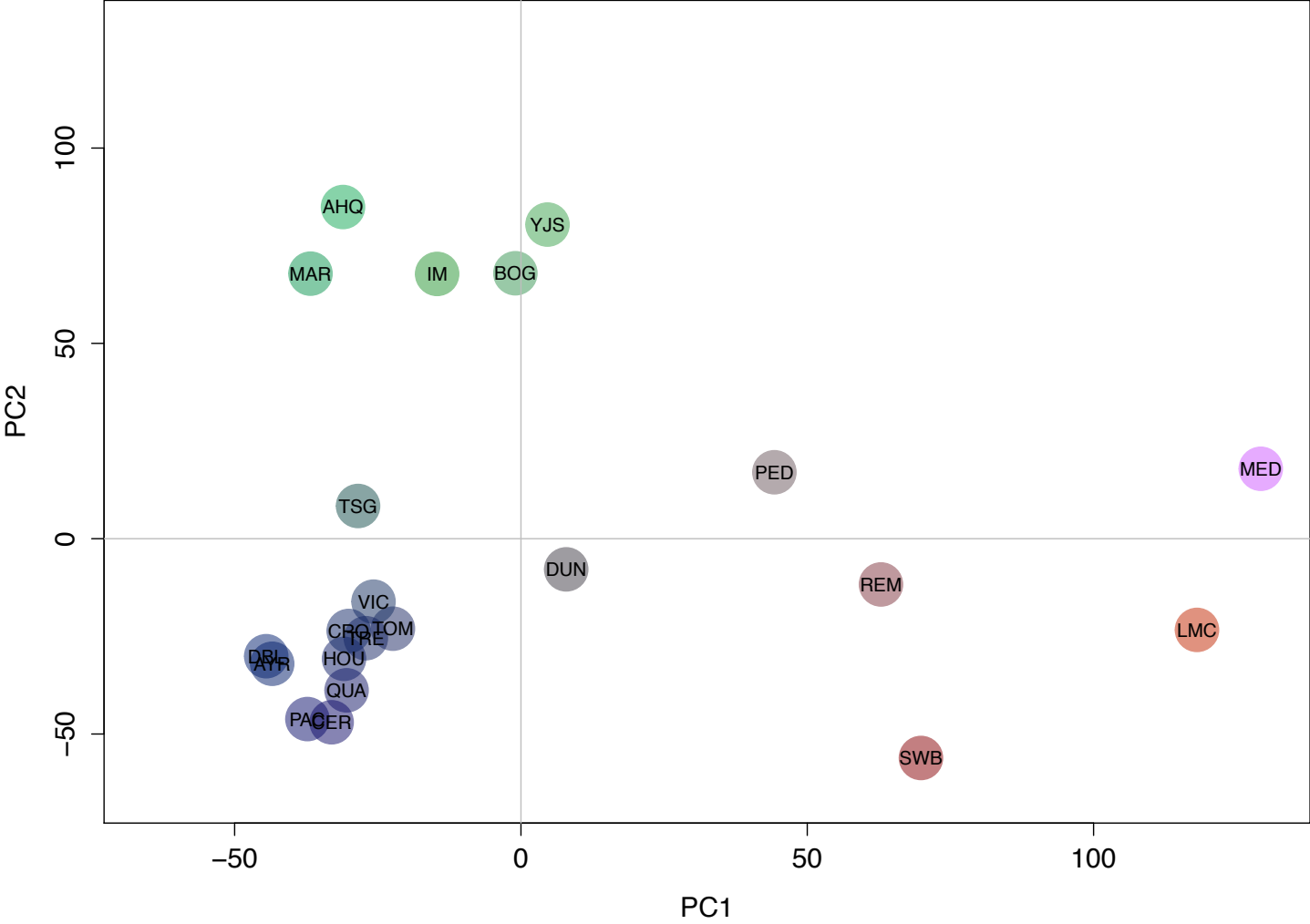
Table S1. Annotated list of 299 genes located within the selective sweeps identified in introduced populations of the yellow monkeyflower, *Mimulus guttatus*, in the United Kingdom. The location of each gene (start and end), and of the selective sweep that contains it, is given in base pairs (bp) with respect to the *M. guttatus* v. 2.0 reference genome (www.phytozome.org).

Table S2. Annotated list of a subset of 28 genes within the selective sweeps described in Table S1 that display reduced synonymous diversity ($\pi_{\text{syn}} < 0.01$).

Supplementary File: Bioinformatics. Commands used for sequence analysis, quality control, and genotyping.







CLR

Linkage Group

